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E.coli

CDS

Escherichia coli K-12 MG1655: b3738

b3738

Entry



BIICLY	D5730 CD5 11,0011
Gene name	atpB, ECK3731, JW3716, papD, uncB
Definition	FO sector of membrane-bound ATP synthase, subunit A (EC:3.6.3.14)
Orthology	KO: K02108 F-type H+-transporting ATPase subunit a [EC:3.6.3.14]
Pathway	PATH: eco00190 Oxidative phosphorylation PATH: eco01100 Metabolic pathways
Class	Metabolism; Energy Metabolism; Oxidative phosphorylation [PATH:eco00190] (BRUTE Merachy)
SSDB	Orthologi (Peralogi) Gene cluster) GFIT
Motif	Pfam: DUF1129 ATP-synt_A PROSITE: ATPASE_A MoNU
Other DBs	RegulonDB: B3738 EcoGene: EG10099 ECCCYC: EG10099 NCBI-GI: 16131606 NCBI-GeneID: 948252 UniProt: POAB98
LinkDB	(All OBs)
Structure	PDB: 1C17 Thumboolis Jmol
Position	complement (39192593920074) (Genome mag)
AA seq	271 aa (AA seg (D8 search) MASENMTPODYIGHHUNNLOLDLRTFSLVDPONPPATFWTINIDSMFFSVVLGLLFLVLF RSVAKKATSGYPCKFQTAIELVIGFVNGSVKDMYHCKSKLIAPLALTIFVWYFLMNLMDL LPIDLLPYIAEHVLGLPALRVVPSADVNVTLSMALGVFILILFYSIKMKGIGGFTKELTL OPPNHWAFIPVNLILEGVSLLSKPVSLGLRLFGNMYAGELIFILIAGLLPWWSQWILNVP WAIFHILIITLQAFIFMVLTIVYLSMASEEH
NT seq	816 nt NI seg +upstream nt +downstream nt atggetteagaaaatatgacgccgcaggattacataggacaccacctgaataaccttcag ctggacctgcqtacattctcggtcggtgcatcacaaaacccccagccaccttctggaca atcaatattgactcatgttcttctggtgtgtgtgtgtctgttgttctgtgtttattccgtagtctgaggtagcgacaaaaagcgaccagcggtgtgccaggtaggt

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